K061129

Attachment B:

Summary of Safety and Effectiveness Prepared in accordance with 21 CFR Part 807.92(c).



GE Healthcare

General Electric Company P.O. Box 414, Milwaukee, WI 53201

Section a):

1.

Submitter: GE Medical Systems, Ultrasound and Primary Care Diagnostics, LLC

PO Box 414

Milwaukee, WI 53201

Contact Person: Patricia Taige.

Safety and Regulatory Engineer

Telephone: 414-721-3222; Fax: 414-721-3854

Date Prepared: April 18, 2006

2.

Device Name: GE LOGIQ 9 Diagnostic Ultrasound BT07,

Ultrasonic Pulsed Echo Imaging System, 21 CFR 892.1560, 90-IYO Ultrasonic Pulsed Doppler Imaging System, 21 CFR 892.1550, 90-IYN

3. Marketed Device: GE LOGIQ 9 Diagnostic Ultrasound System K011188/K030934/K032656/K040251

A device currently in commercial distribution.

- 4. <u>Device Description</u>: The GE LOGIQ 9 is a full-featured general-purpose diagnostic ultrasound system. It consists of a mobile console approximately 64 cm wide, 90 cm deep and 140-160 cm (adjustable) high that provides digital acquisition, processing and display capability. The user interface includes a computer keyboard, specialized controls and a color video CRT or LCD display and LCD touch panel. This modification will provide users with an intima media thickness measurement tool and enhanced 3D image manipulation.
- 5. Indications for Use: The device is intended for use by a qualified physician for u'trasound evaluation of Fetal; Abdominal; Pediatric; Small Organ (breast, testes, thyroid); Neonatal Cephalic; Adult Cephalic; Cardiac (adult and pediatric); Peripheral Vascular; Musculo-skeletal Conventional and Superficial; Urology (including prostate); Transrectal; Transvaginal; and Intraoperative (abdominal, thoracic, vascular and neurosurgical).
- 6. Comparison with Predicate Device: The GE LOGIQ 9 BT07 is of a comparable type and substantially equivalent to the current GE LOGIQ 9 and GE LOGIQ 7. It has the same technological characteristics, key safety and effectiveness features, physical design, construction, and materials, and has the same intended uses and basic operating modes as the predicate device.

Section b):

- 1. Non-clinical Tests: The device has been evaluated for acoustic output, biocompatibility, cleaning and disinfection effectiveness as well as thermal, electrical and mechanical safety, and has been found to conform with applicable medical device safety standards.
- 2. Clinical Tests: None required.
- 3. Conclusion: Intended uses and other key features are consistent with traditional clinical practice, FDA guidelines, and established methods of patient examination. The design and development process of the manufacturer conforms with 21 CFR 820, ISO 9001:2000 and ISO 13485 quality systems. The device conforms to applicable medical device safety standards and compliance is verified through independent evaluation with ongoing factory surveillance. Diagnostic ultrasound has accumulated a long history of safe and effective performance. Therefore, it is the opinion of GE Medical Systems that the GE LOGÍQ 9 BT07 Diagnostic Ultrasound is substantially equivalent with respect to safety and effectiveness to devices currently cleared for market.



Food and Drug Administration 9200 Corporate Boulevard Rockville MD 20850

JUN = 1 2006

Ms. Patricia Taige
Safety & Regulatory Engineer
General Electric Company
GE Medical Systems, Ultrasound and Primary Care Diagnostics, LLC
99000 Innovation Drive
WAUWATOSA WI 53226

Re: K061129

Trade Name: GE LOGIQ 9 Ultrasound System

Regulation Number: 21 CFR 892.1550

Regulation Name: Ultrasonic pulsed doppler imaging system

Regulation Number: 21 CFR 892.1560

Regulation Name: Ultrasonic pulsed echo imaging system

Regulation Number: 21 CFR 892.1570

Regulation Name: Diagnostic ultrasonic transducer

Regulatory Class: II

Product Code: IYN, IYO, and ITX

Dated: April 19, 2006 Received: April 24, 2006

Dear Ms. Taige:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and we have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration.

This determination of substantial equivalence applies to the following transducers intended for use with the GE LOGIQ 9 Ultrasound System, as described in your premarket notification:



Transducer Model Number

4D8C 4D16L 4C 9L i739 or t739 P5D

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to such additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

This determination of substantial equivalence is granted on the condition that prior to shipping the first device, you submit a postclearance special report. This report should contain complete information, including acoustic output measurements based on production line devices, requested in Appendix G, (enclosed) of the Center's September 30, 1997 "Information for Manufacturers Seeking Marketing Clearance of Diagnostic Ultrasound Systems and Transducers." If the special report is incomplete or contains unacceptable values (e.g., acoustic output greater than approved levels), then the 510(k) clearance may not apply to the production units which as a result may be considered adulterated or misbranded.

The special report should reference the manufacturer's 510(k) number. It should be clearly and prominently marked "ADD-TO-FILE" and should be submitted in duplicate to:

Food and Drug Administration Center for Devices and Radiological Health Document Mail Center (HFZ-401) 9200 Corporate Boulevard Rockville, Maryland 20850

This letter will allow you to begin marketing your device as described in your premarket notification. The FDA finding of substantial equivalence of your device to a legally marketed predicate device results in a classification for your device and thus permits your device to proceed to market.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please contact the Office of Compliance at (240) 276-0120. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21 CFR Part 807.97). You may obtain other general information on your responsibilities under the Act from the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (301) 443-6597 or at its Internet address http://www.fda.gov/cdrh/industry/support/index.html

If you have any questions regarding the content of this letter, please contact Ralph Shuping at (301) 594-1212.

Sincerely yours,

Davidh. Lynnfor Nancy C. Brogdon

Director, Division of Reproductive, Abdominal and Radiological Devices

Office of Device Evaluation

Center for Devices and Radiological Health

Enclosure

GE LOGIQ 9 Ultrasound System

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Mode of Operation										
Clinical Application Anatomy/Region of Interest	В .	М	PW Doppler	CW Doppler	Color Doppler	Color M Doppler	Power Doppler		Harmonic Imaging	Coded Pulse	Other
Ophthalmic											
Fetal / Obstetrics	Р	Р	Р	Р	Р	Р	Р	Р	Р	Р	
Abdominal ^[1]	Р	Р	Р	Р	Р	Р	Р	Р	P	Р	
Pediatric	P	Р	Р	Р	Р	Р	Р	Р	Р	Р	
Small Organ ^[2]	Р	Р	Р		Р	P	Р	Р	P	Р	
Neonatal Cephalic	Р	Р	Р	Р	Р	Р	Р	P __	Р	Р	
Adult Cephalic	Р	Р	Р	Р	Р	P	Р	Р	P	·Р	
Cardiac ^[3]	Р	Р	Р	P	Р	Р	Р	Р	P	Р	
Peripheral Vascular	Р	P	Р	P	p	Р	Р	Р	Р	Р	
Musculo-skeletal Conventional	Р	Р	Р		Р	Р	Р	Р	P	Р	
Musculo-skeletal Superficial	Р	Р	Р		Р	Р	Р	Р	Р	Р	
Other ^[4]	Р	Р	Р	Р	Р	Р	Р	Р	Р	P	
Exam Type, Means of Access											
Transesophageal				_							
Transrectal	Р	Р	Р		Р	Р	Р	Р	Р	Р	
Transvaginal	Р	Р	Р		Р	Р	Р	P	P	Р	ļ
Transuretheral									***		
Intraoperative ⁽⁵⁾	P	Р	Р		Р	Р	Р	Р	Р	Р	
Intraoperative Neurological	Р	P	P		Р	Р	Р	P	Р	Р	
Intravascular											
Laparoscopic											

N = new indication; P = previously cleared by FDA; E = added under Appendix E

N	lotes:	[1]	Abdominal	includes	renal,	GYN/P	elvic
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- [2] Small organ includes breast, testes, thyroid.
- [3] Cardiac is Adult and Pediatric.
- [4] Other use includes Urology/Prostate
- [5] Intraoperative includes abdominal, thoracic (cardiac), and vascular (PV).
- [*] Combined modes are B/M, B/Color M, B/PWD or CWD, B/Color/PWD or CWD, B/Power/PWD.

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 Concurrence of CDRH, Office of Device Evaluation (ODE)	

(Division Sign-Off)

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Division of Reproductive, Abdominal,

and Radiological Devices 510(k) Number

K061129

GE LOGIQ 9 with 4D8C Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

					Mode	of Ope	eration				
Clinical Application Anatomy/Region of Interest	В	М	PW Doppler	CW Doppler	Color Doppler	Color M Doppler	Power Doppler		Harmonic Imaging	Coded Pulse	Other
Ophthalmic											
Fetal / Obstetrics	Ε	E	E_		E	E	E	E	Е	E	
Abdominal ^[1]	E	E	Ε		E	E	E	E	E	E	<u> </u>
Pediatric	E	E	E		E	E	Ε	E	E	E	
Small Organ ^[2]	E	E	E		E	E	E	E	E	E	ļ
Neonatal Cephalic	E	E	E		E	E	E	E.	Ε	E	
Adult Cephalic											ļ
Cardiac ^[3]											
Peripheral Vascular		ļ <u>-</u>									<u> </u>
Musculo-skeletal Conventional						<u></u>					<u> </u>
Musculo-skeletal Superficial								ļ			
Other ^[4]											
Exam Type, Means of Access					<u> </u>			<u> </u>			
Transesophageal		ļ		ļ <u> </u>							
Transrectal											 -
Transvaginal		<u> </u>							ļ <u>-</u>		
Transuretheral			<u> </u>								
Intraoperative ^[5]		<u> </u>	ļ			ļ	ļ	ļ	1		
Intraoperative Neurological							<u> </u>	<u> </u>		-	
Intravascular					ļ		<u> </u>			<u> </u>	-
Laparoscopic			<u> </u>				<u> </u>	<u> </u>			<u></u>

N = new indication; P = previously cleared by FDA; E = added under Appendix E

Notes:	[1]	Abdomina	Lincludes	renal,	GYN/Pelvic
--------	-----	----------	-----------	--------	------------

- [2] Small organ includes breast, testes, thyroid.
- [3] Cardiac is Adult and Pediatric.
- [4] Other use includes Urology/Prostate
- [5] Intraoperative includes abdominal, thoracic (cardiac), and vascular (PV).
- [*] Combined modes are B/M, B/Color M, B/PWD or CWD, B/Color/PWD or CWD, B/Power/PWD.

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Division of Reproductive, Abdominal,

and Radiological Devices 510(k) Number _____

al Devices **K06/12**

GE LOGIQ 9 with 4D16L Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

		Mode of Operation									
Clinical Application Anatomy/Region of Interest	В	М	PW Doppler	CW Doppler	Color Doppler	Color M Doppler	T	Combined	Harmonic Imaging	Coded Pulse	Other
Ophthalmic											
Fetal / Obstetrics											
Abdominal ^[1]	Е	E	E		E		E	E	E	E	
Pediatric	E	E	E		E		E	E	E	E	
Small Organ ^[2]	E	E	E		E		E	E	E	Ε	
Neonatal Cephalic	E	E	E		E		E	E,	E	Е	
Adult Cephalic											
Cardiac ^[3]											
Peripheral Vascular	Е	Е	Е		E		Е	Е	E	Е	
Musculo-skeletal Conventional	Е	Е	E		E		Е	Ε	E	ш	
Musculo-skeletal Superficial	E	E	E		E		E	E	E	Ε	
Other ^[4]											ļ
Exam Type, Means of Access											
Transesophageal											
Transrectal											
Transvaginal											
Transuretheral											
Intraoperative ^[5]											
Intraoperative Neurological											
Intravascular											
Laparoscopic											

N = new indication; P = previously cleared by FDA; E = added under Appendix E

Notes:	[1]	Abdominal	includes	renal	GYN/Pelvic
	1	riodomina	moraco	I CI ICII,	CHITTI CIVIC

- [2] Small organ includes breast, testes, thyroid.
- [3] Cardiac is Adult and Pediatric.
- [4] Other use includes Urology/Prostate
- [5] Intraoperative includes abdominal, thoracic (cardiac), and vascular (PV).
- [*] Combined modes are B/M, B/Color M, B/PWD or CWD, B/Color/PWD or CWD, B/Power/PWD.

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GE LOGIO 9 with 4C Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Mode of Operation										
Clinical Application Anatomy/Region of Interest	В	М	PW Doppler	CW Doppler	Color Doppler	Color M Doppler		Combined Modes	Harmonic Imaging	Coded Pulse	Other
Ophthalmic											
Fetal / Obstetrics	E	E	Е		E	Ε	E	E	E	E	
Abdominal ⁽¹⁾	Ε	E	E		E	E	E	Ε	E	E	
Pediatric											
Small Organ ^[2]											
Neonatal Cephalic											
Adult Cephalic											
Cardiac ^[3]											
Peripheral Vascular											
Musculo-skeletal Conventional	Е	E	Ε		E	E	E	E	Ε	E	
Musculo-skeletal Superficial											
Other ^[4]	Ε	E	E		E	E	E	E	Е	E	
Exam Type, Means of Access											
Transesophageal											
Transrectal											
Transvaginal											
Transuretheral											
Intraoperative ^[5]									ļ.,,,		
Intraoperative Neurological											ļ
Intravascular											<u> </u>
Laparoscopic											

N = new indication; P = previously cleared by FDA; E = added under Appendix E

Notes:	[1]	Abdominal	Lincludes	renal,	GYN/Pelvic
--------	-----	-----------	-----------	--------	------------

- [2] Small organ includes breast, testes, thyroid.
- [3] Cardiac is Adult and Pediatric.
- [4] Other use includes Urology/Prostate
- [5] Intraoperative includes abdominal, thoracic (cardiac), and vascular (PV).
- [*] Combined modes are B/M, B/Color M, B/PWD or CWD, B/Color/PWD or CWD, B/Power/PWD.

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Division of Reproductive, Abdominal,

and Radiological Devices

510(k) Number _____

GE LOGIQ 9 with 9L Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

					Mode	of Ope	eration				
Clinical Application Anatomy/Region of Interest	В	М	PW Doppler	CW Doppler	Color Doppler	Color M Doppler	Power Doppler	Combined Modes	Harmonic Imaging	Coded Pulse	Other
Ophthalmic											
Fetal / Obstetrics	E	E	E		E		E	E	Ε	E	
Abdominal ^[1]	Ε	E	E		E		E	Ε	Е	Ε	
Pediatric											
Small Organ ^[2]	£	E	Ε		E	<u>.</u>	Е	E	Ε	Ε	
Neonatal Cephalic											
Adult Cephalic											
Cardiac ^[3]											
Peripheral Vascular	Е	E	E		E		Ε	E.	E	Ε	
Musculo-skeletal Conventional	Ε	E	E		E		E	E	E	E	
Musculo-skeletal Superficial	E	E	Ε		Ε		Ε	Ε	E	E	
Other ^[4]											
Exam Type, Means of Access											
Transesophageal								:			
Transrectal											
Transvaginal										· 	
Transuretheral											
Intraoperative ^[5]											
Intraoperative Neurological										•	
Intravascular		_									
Laparoscopic			-								

N = new indication; P = previously cleared by FDA; E = added under Appendix E

Notes: [1] Abdominal includes	s renal GYN/Pelvi	C.
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- [2] Small organ includes breast, testes, thyroid.
- [3] Cardiac is Adult and Pediatric.
- [4] Other use includes Urology/Prostate
- [5] Intraoperative includes abdominal, thoracic (cardiac), and vascular (PV).
- [*] Combined modes are B/M, B/Color M, B/PWD or CWD, B/Color/PWD or CWD, B/Power/PWD.

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Division of Reproductive, Abdominal, and Radiological Devices
510(k) Number

Diagnostic Ultrasound Indications for Use Form GE LOGIQ 9 with i739 or t739 Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

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			1	T	1	of Ope	T		г		
Clinical Application	В	М	PW Doppler	CW Doppler	Color Doppler	Color M Doppler	Power Doppler	Combined Modes	Harmonic Imaging	Coded Pulse	Other
Anatomy/Region of Interest			Dobbiei	Dopplei	Dobbiei	Dopplet	Doppier	Wodes	imaging	1 4,50	
Ophthalmic											
Fetal / Obstetrics				ļ							
Abdominal ^[1]			<u> </u>	ļ							ļ
Pediatric											<u> </u>
Small Organ ^[2]											ļ
Neonatal Cephalic											<u> </u>
Adult Cephalic						ļ					
Cardiac ^[3]										<u> </u>	<u> </u>
Peripheral Vascular							ļ			:	<u> </u>
Musculo-skeletal Conventional											
Musculo-skeletal Superficial											
Other ^[4]											ļ
Exam Type, Means of Access											<u> </u>
Transesophageal											<u> </u>
Transrectal			<u> </u>							ļ <u>-</u>	-
Transvaginal											
Transuretheral											ļ
Intraoperative ^[5]	E		Ε		E	E	E	E	E	E	ļ
Intraoperative Neurological	Е		E		E	Е	E	E	E	E	
Intravascular											ļ
Laparoscopic			1		1						

N = new indication; P = previously cleared by FDA; E = added under Appendix E

Notes:	[1]	Abdomir	al incl	ludes	renal	GYN/F	Pelvic
MURGO.	111	AUGUITHI	iai ii ci	เนนธอ	i Cilai.	O DIVI	CIVIL.

- [2] Small organ includes breast, testes, thyroid.
- [3] Cardiac is Adult and Pediatric.
- [4] Other use includes Urology/Prostate
- [5] Intraoperative includes abdominal, thoracic (cardiac), and vascular (PV).
- [*] Combined modes are B/M, B/Color M, B/PWD or CWD, B/Color/PWD or CWD, B/Power/PWD.

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(Division Sign-Off) Division of Reproductive, Abdominal,

and Radiological Devices Ko6/1/24

510(k) Number ___

GE LOGIQ 9 with P5D Transducer

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Mode of Operation										
Clinical Application Anatomy/Region of Interest	В	М	PW Doppler	CW Doppler	Color Doppler	Color M Doppler		Combined Modes	Harmonic Imaging	Coded Pulse	Other
Ophthalmic											
Fetal / Obstetrics											
Abdominal ^[1]											
Pediatric											
Small Organ ^[2]									ļ		ļ
Neonatal Cephalic											
Adult Cephalic											
Cardiac ^[3]						ļ 	ļ				ļ
Peripheral Vascular				E							
Musculo-skeletal Conventional									ļ		ļ
Musculo-skeletal Superficial		ļ									ļ
Other ^[4]		ļ						ļ			<u></u>
Exam Type, Means of Access		ļ						ļ			ļ
Transesophageal				ļ <u>.</u>	<u> </u>			ļ			-
Transrectal		ļ							ļ		ļ <u> </u>
Transvaginal											ļ
Transuretheral		<u> </u>						<u> </u>			
Intraoperative ^[5]								ļ	-	ļ	
Intraoperative Neurological			_						ļ		
Intravascular											ļ
Laparoscopic					<u> </u>	<u> </u>		<u></u>	<u> </u>	<u> </u>	<u> </u>

N = new indication; P = previously cleared by FDA; E = added under Appendix E

Notes:	[1]	Abdominal	includes	renal	GYN/Pelvio
noies.	111	Abuonniai	Includes	Tenal.	CHINECING

- [2] Small organ includes breast, testes, thyroid.
- [3] Cardiac is Adult and Pediatric.
- [4] Other use includes Urology/Prostate
- (FV).

1 Combined mod	es are B/M, B/Color M, B/PWD or CWD, B/Color/PWD or CWD, B/Power/PV
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	Concurrence of CDRH, Office of Device Evaluation (ODE)
	David A. begen
	(Division Sign-Off)
	Division of Reproductive, Abdominal,
	and Radiological Devices KO61129

510(k) Number _____